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ANSWER 1 OF 3 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 1
L15
     2004:182819 CAPLUS
AN
DN
     140:237530
     Processes and catalysts for the preparation of 2-chloro-1,1,1,2,3,3,3-
TI
     heptafluoropropane, hexafluoropropene and 1,1,1,2,3,3,3-heptafluoropropane
     Nappa, Mario J.; Rao, Velliyur Nott Mallikarjuna; Rosenfeld, H. David;
IN
     Subramoney, Shekhar; Subramanian, Munirpallam A.; Sievert, Allen C.
     E.I. du Pont de Nemours and Company, USA
PA
SO
     PCT Int. Appl., 29 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LΑ
FAN.CNT 1
                                                                   DATE
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                                DATE
                                          APPLICATION NO.
     PATENT NO.
                                ______
     _____
                         _ - - -
                                          WO 2003-US26331
                                20040304
                                                                   20030821
     WO 2004018397
                         A1
PI
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
             PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
             TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
             CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC,
             NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ,
             GW, ML, MR, NE, SN, TD, TG
                                20020822
PRAI US 2002-405222P
                          Ρ
     CASREACT 140:237530; MARPAT 140:237530
OS
     A process for the preparation of 2-chloro-1,1,1,3,3,3-heptafluoropropane is
AΒ
     described which involves: (a) contacting a mixture comprising hydrogen
     fluoride, chlorine, and at least one starting material selected from
     halopropenes CX3CCl:CX2 (X = F, Cl; Y = H, Cl, F; provided that the number of
     X and Y which are F totals ≤6) and halopropanes CX3CClYCX3, where
     each with a chlorofluorination catalyst in a reaction zone to produce a
     product mixture comprising CF3CClFCF3, HCl, HF, and underfluorinated
     halogenated hydrocarbon intermediates. The chlorofluorination catalyst
     comprises at least one chromium-containing component selected from (i) a
     crystalline alpha-chromium oxide where at least 0.05 atom% of the chromium
     atoms in the alpha-chromium oxide lattice are replaced by nickel,
     trivalent cobalt or both nickel and trivalent cobalt, provided that no
     more than 2 atom% of the chromium atoms in the alpha-chromium oxide
     lattice are replaced by nickel and that the total amount of chromium atoms
     in the alpha-chromium oxide lattice that are replaced by nickel and
     trivalent cobalt is no more than 6 atom% , and (ii) a fluorinated crystalline
     oxide of (i). Also described is a process for the manufacture of a mixture of
     HFC-227ea and hexafluoropropene by reacting a starting mixture comprising
     CFC-217ba and hydrogen in the vapor phase at an elevated temperature,
optionally
     in the presence of a hydrogenation catalyst.
              THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 2
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
L15 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 2
AN
     1992:20676 CAPLUS
     116:20676
DN
     Multistep synthesis of hexafluoropropylene from propane and propylene
TI
     Webster, James Lang; McCann, Elrey Lorne; Bruhnke, Douglas William; Lerou,
IN
     Jan Joseph; Manogue, William Henry; Manzer, Leo Ernest; Swearingen, Steven
     Henry; Trofimenko, Swiatoslaw; Bonifaz, Cristobal
     du Pont de Nemours, E. I., and Co., USA
PΑ
     Eur. Pat. Appl., 33 pp.
SO
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CODEN: EPXXDW

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DT
         Patent
LA
         English
FAN.CNT 1
                                KIND DATE
                                                                           APPLICATION NO.
                                                                                                                  DATE
         PATENT NO.
                                                                           _ _ _ _
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                                                       19910626
                                                                           EP 1990-313951
                                                                                                                 19901219
                                            A1
PI
         EP 434409
                                          B1
                                                       19941012
         EP 434409
R: DE, FR, GB, IT

US 5057634

CA 2032273

CA 2032273

CA 2032273

CA 2298099

CA 2298099

CA 2298099

CB 20020108

CB 200
               R: DE, FR, GB, IT
                                                                           US 1989-452402
                                                                                                                 19891219
                                                                           CA 1990-2032273
                                                                                                                   19901214
                                                                           CA 1990-2298099
                                                                                                                19901214
                                                                                                                   19901219
                                                                           JP 1990-411690
         Hexafluoropropylene (I) is prepared by (1) chlorofluorination of at least
AΒ
         one of propane, propylene, and partially halogenated C3 acyclic
         hydrocarbons with HF and Cl in the presence of a chlorofluorination
         catalyst to produce CF3CFClCF3 (II) and other chlorofluorocarbons such as
         C3F4Cl4, C3H5Cl3, CF3CFClCF2Cl, CF3CCl2CF3, and CF3CCl2CCl3 which are
         mostly recyclable to the same chlorofluorination step to give II and (2)
         dehalogenation of II to form I in the presence of a CuO-NiO-Cr2O3-CaF2
         (and-MoO3) catalyst containing at least one of K, Cs, or Rb. In this process
         there is substantially no perfluoroisobutylene produced as a byproduct
         which is extremely toxic and is costly to remove and destroy. Thus,
         Cr203.3H2O was charged to an Inconel tubular reactor and treated with a
         flow of HF at 400° for dehydration and thereto HF 90, Cl 35, and
         propylene 1.5 mol/h were fed at 440° and 790 kPa to give II 75,
         C3F6Cl2 7, C3F5Cl3 5, C3F7H 3, C3F6ClH 5, C3F8 2 and C2F5Cl 2%.
         (mol) mixture of H and a II feed containing II 79, CF3CF2CF2Cl 17, and
         CF3CCl:CF2 0.7% was passed over a catalyst CuO/NiO/Cr2O3/2.7 CaF2 containing
         7.9 weight% K at 402° to give 97% I with 63% conversion of II.
         ANSWER 3 OF 3 USPATFULL on STN
L15
            91:84577 USPATFULL
ΑN
            Multistep synthesis of hexafluoropropylene
TI
            Webster, James L., Parkersburg, WV, United States
 IN
            McCann, Elrey L., Mendenhall, PA, United States
            Bruhnke, Douglas W., Landenberg, PA, United States
            Lerou, Jan J., Chadds Ford, PA, United States
            Manogue, William H., Newark, DE, United States
            Manzer, Leo E., Wilmington, DE, United States
            Swearingen, Steven H., Wilmington, DE, United States
            Trofimenko, Swiatoslaw, Wilmington, DE, United States
            Bonifaz, Cristobal, Conway, MA, United States
            E. I. Du Pont de Nemours and Company, Wilmington, DE, United States
PΑ
             (U.S. corporation)
            US 5057634
                                                    19911015
PΙ
            US 1989-452402
                                                   19891219 (7)
AΙ
DT
            Utility
FS
            Granted
EXNAM Primary Examiner: Siegel, Alan
            Huntley, Donald W.
LREP
            Number of Claims: 6
CLMN
            Exemplary Claim: 1
ECL
            3 Drawing Figure(s); 3 Drawing Page(s)
DRWN
LN.CNT 1458
 CAS INDEXING IS AVAILABLE FOR THIS PATENT
            The present invention relates to multistep syntheses of
AΒ
            hexafluoropropylene from acyclic three-carbon hydrocarbons or partially
            halogenated acyclic three-carbon hydrocarbons. In all these syntheses
            the first step is a vapor-phase chlorofluorination of the starting
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material to one or more saturated chlorofluorocarbons. Novel catalysts

are also provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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2004:182753 CAPLUS
AN
DN
     140:201451
     Cobalt-substituted chromium oxide compositions, their preparation, and
TI
     their use as catalysts and catalyst precursors
    Nappa, Mario J.; Rao, Velliyur Nott Mallikarjuna; Rosenfeld, David H.;
IN
     Subramoney, Shekhar; Subramanian, Munirpallam A.; Sievert, Allen C.
     E.I. du Pont de Nemours and Company, USA
PA
     PCT Int. Appl., 68 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
                        KIND DATE APPLICATION NO. DATE
     PATENT NO.
                               _____
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                               20040304 WO 2003-US26326
                                                                 20030821
                        A2
    WO 2004018093
PI
                         A3 20040422
     WO 2004018093
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
             PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
             TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
             CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC,
             NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ,
             GW, ML, MR, NE, SN, TD, TG
PRAI US 2002-405220P
                        P 20020822
     A crystalline \alpha\text{-chromium} oxide where 0.05-6 atom% of the chromium atoms
     in the \alpha-chromium oxide lattice are replaced by trivalent cobalt
     (Co+3) atoms is disclosed. Also disclosed is a chromium-containing catalyst
     composition comprising as a chromium-containing component the crystalline
     cobalt-substituted \alpha-chromium oxide; and a method for preparing a
     composition comprising the crystalline cobalt-substituted \alpha-chromium oxide.
     The method involves (a) co-precipitating a solid by adding ammonium hydroxide
to
     an aqueous solution of a soluble cobalt salt and a soluble trivalent chromium
salt that
     contains ≥3 mol of nitrate/mol of chromium in the solution and has a
     cobalt concentration 0.05-6 mol% of the total concentration of cobalt and
chromium in
     the solution; and after at least three moles of ammonium per mol of chromium
     in the solution has been added to the solution, (b) collecting the
co-precipitated
     solid formed in (a); (c) drying the collected solid; and (d) calcining the
     dried solid. Also disclosed is a chromium-containing catalyst composition
     comprising a chromium-containing component prepared by treating the crystalline
     cobalt-substituted -chromium oxide with a fluorinating agent; and a
     process for changing the fluorine distribution (i.e., content and/or
     arrangement) in a hydrocarbon or halogenated hydrocarbon in the presence
     of a catalyst. The process involves using as the catalyst a composition
     comprising the crystalline cobalt-substituted \alpha-chromium oxide and/or the
     treated cobalt-substituted \alpha-chromium oxide.
L18 ANSWER 2 OF 12 USPATFULL on STN
       2004:134127 USPATFULL
AN
       Materials and methods for the production and purification of
TI
       chlorofluorocarbons and hydrofluorocarbons
       Iikubo, Yuichi, West Lafayette, IN, UNITED STATES
IN
       Owens, Stephen, White Pine, TN, UNITED STATES
       Cohn, Mitchel, West Lafayette, IN, UNITED STATES
```

Brandstadter, Stephan M., Indianapolis, IN, UNITED STATES

ANSWER 1 OF 12 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 1

L18

Hedrick, Vicki E., Brookston, IN, UNITED STATES Boggs, Janet K., Brownsburg, IN, UNITED STATES Qian, John, West Lafayette, IN, UNITED STATES Sacarias, Julie, El Dorado, AR, UNITED STATES 20040527 US 2004102664 **A**1 PΙ US 2003-699491) 20031031 (10) Α1 AIRLIDTUtility

Division of Ser. No. US 2001-966158, filed on 28 Sep 2001, PENDING

FS APPLICATION

WELLS ST. JOHN P.S., 601 W. FIRST AVENUE, SUITE 1300, SPOKANE, WA, 99201 LREP

Number of Claims: 88 CLMNExemplary Claim: 1 ECL 7 Drawing Page(s) DRWN

LN.CNT 1969

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Methods and materials are provided for the production of essentially isomerically pure perhalogenated and partially halogenated compounds. One embodiment of the present invention provides a process for the production of essentially isomerically pure CFC-216aa. Other embodiments include processes for the production of CFC-217ba and HFC-227ea. Particular embodiments of the present invention provide separation techniques for the separation of chlorofluorocarbons from HF, from other chlorofluorocarbons, and the separation of isomers of halogenated compounds. Still other embodiments of the present invention provide catalytic synthetic techniques that demonstrate extended catalyst lifetime. In other embodiments, the present invention provides catalytic techniques for the purification of isomeric mixtures.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 3 OF 12 USPATFULL on STN L18

2004:134126 USPATFULL AN

Materials and methods for the production and purification of TIchlorofluorocarbons and hydrofluorocarbons

Iikubo, Yuichi, West Lafayette, IN, UNITED STATES INOwens, Stephen, White Pine, TN, UNITED STATES Cohn, Mitchel, West Lafayette, IN, UNITED STATES Brandstadter, Stephan M., Indianapolis, IN, UNITED STATES Hedrick, Vicki E., Brookston, IN, UNITED STATES Boggs, Janet K., Brownsburg, IN, UNITED STATES Qian, John, West Lafayette, IN, UNITED STATES Sacarias, Julie, El Dorado, AR, UNITED STATES

US 2004102663 A1 20040527 PIUS 2003-698923 20031031 (10) **A**1

ΑI Division of Ser. No. US 2001-966158, filed on 28 Sep 2001, PENDING RLI

DTUtility

APPLICATION FS

WELLS ST. JOHN P.S., 601 W. FIRST AVENUE, SUITE 1300, SPOKANE, WA, 99201 LREP

Number of Claims: 88 CLMN Exemplary Claim: 1 ECL 7 Drawing Page(s) DRWN

LN.CNT 1970

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Methods and materials are provided for the production of essentially ABisomerically pure perhalogenated and partially halogenated compounds. One embodiment of the present invention provides a process for the production of essentially isomerically pure CFC-216aa. Other embodiments include processes for the production of CFC-217ba and HFC-227ea. Particular embodiments of the present invention provide separation techniques for the separation of chlorofluorocarbons from HF, from other chlorofluorocarbons, and the separation of isomers of halogenated compounds. Still other embodiments of the present invention provide catalytic synthetic techniques that demonstrate extended catalyst lifetime. In other embodiments, the present invention provides catalytic techniques for the purification of isomeric mixtures.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

LN.CNT 1967

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L18 ANSWER 4 OF 12 USPATFULL on STN
       2004:134125 USPATFULL
AN
       Processes for purifying chlorofluorinated compounds
{
m TI}
       Iikubo, Yuichi, West Lafayette, IN, UNITED STATES
IN
       Owens, Stephen, White Pine, TN, UNITED STATES
       Cohn, Mitchel, West Lafayette, IN, UNITED STATES
       Brandstadter, Stephan M., Indianapolis, IN, UNITED STATES
       Hedrick, Vicki E., Brookston, IN, UNITED STATES
       Boggs, Janet K., Brownsburg, IN, UNITED STATES
       Qian, John, West Lafayette, IN, UNITED STATES
       Sacarias, Julie, El Dorado, AR, UNITED STATES
       US 2004102662 A1 20040527
US 2003-698730 A1 20031031
PI
                               20031031 (10)
ΑI
       Division of Ser. No. US 2001-966158, filed on 28 Sep 2001, PENDING
RLI
       Utility
DT
       APPLICATION
FS
       WELLS ST. JOHN P.S., 601 W. FIRST AVENUE, SUITE 1300, SPOKANE, WA, 99201
LREP
       Number of Claims: 88
CLMN
       Exemplary Claim: 1
ECL
       7 Drawing Page(s)
DRWN
LN.CNT 1974
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Methods and materials are provided for the production of essentially
AB
       isomerically pure perhalogenated and partially halogenated compounds.
       One embodiment of the present invention provides a process for the
       production of essentially isomerically pure CFC-216aa. Other embodiments
       include processes for the production of CFC-217ba and HFC-227ea.
       Particular embodiments of the present invention provide separation
       techniques for the separation of chlorofluorocarbons from HF, from other
       chlorofluorocarbons, and the separation of isomers of halogenated
       compounds. Still other embodiments of the present invention provide
       catalytic synthetic techniques that demonstrate extended catalyst
       lifetime. In other embodiments, the present invention provides catalytic
       techniques for the purification of isomeric mixtures.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
    ANSWER 5 OF 12 USPATFULL on STN
L18
       2004:134124 USPATFULL
AN
       Processes for purifying chlorofluorinated compounds and processes for
TI
       purifying CF3CFHCF3
       Iikubo, Yuichi, West Lafayette, IN, UNITED STATES
IN
       Owens, Stephen, White Pine, TN, UNITED STATES
       Cohn, Mitchel, West Lafayette, IN, UNITED STATES
       Brandstadter, Stephan M., Indianapolis, IN, UNITED STATES
       Hedrick, Vicki E., Brookston, IN, UNITED STATES
       Boggs, Janet K., Brownsburg, IN, UNITED STATES
       Qian, John, West Lafayette, IN, UNITED STATES
       Sacarias, Julie, El Dorado, AR, UNITED STATES
                          A1
                               20040527
       US 2004102661
PI
       US 2003-698579
                               20031031 (10)
                         A1
AI
       Division of Ser. No. US 2001-966158, filed on 28 Sep 2001, PENDING
\mathtt{RLI}
       Utility
DT
FS
       APPLICATION
                            601 W. FIRST AVENUE, SUITE 1300, SPOKANE, WA,
       WELLS ST. JOHN P.S.,
LREP
       Number of Claims: 88
CLMN
       Exemplary Claim: 1
ECL
       7 Drawing Page(s)
DRWN
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AB Methods and materials are provided for the production of essentially isomerically pure perhalogenated and partially halogenated compounds. One embodiment of the present invention provides a process for the production of essentially isomerically pure CFC-216aa. Other embodiments include processes for the production of CFC-217ba and HFC-227ea. Particular embodiments of the present invention provide separation techniques for the separation of chlorofluorocarbons from HF, from other chlorofluorocarbons, and the separation of isomers of halogenated compounds. Still other embodiments of the present invention provide catalytic synthetic techniques that demonstrate extended catalyst lifetime. In other embodiments, the present invention provides catalytic techniques for the purification of isomeric mixtures.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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ANSWER 6 OF 12 USPATFULL on STN
L18
AN
       2004:134123 USPATFULL
TI
       Processes for halogenating compounds
       Iikubo, Yuichi, West Lafayette, IN, UNITED STATES
IN
       Owens, Stephen, White Pine, TN, UNITED STATES
       Cohn, Mitchel, West Lafayette, IN, UNITED STATES
       Brandstadter, Stephan M., Indianapolis, IN, UNITED STATES
       Hedrick, Vicki E., Brookston, IN, UNITED STATES
       Boggs, Janet K., Brownsburg, IN, UNITED STATES
       Qian, John, West Lafayette, IN, UNITED STATES
       Sacarias, Julie, El Dorado, AR, UNITED STATES
                               20040527
PI
       US 2004102660
                         Α1
       US 2003-698731
ΑI
                        A1
                               20031031 (10)
       Division of Ser. No. US 2001-966158, filed on 28 Sep 2001, PENDING
RLI
       Utility
DT
FS
       APPLICATION
       WELLS ST. JOHN P.S., 601 W. FIRST AVENUE, SUITE 1300, SPOKANE, WA, 99201
LREP
CLMN
       Number of Claims: 88
ECL
       Exemplary Claim: 1
       7 Drawing Page(s)
DRWN
LN.CNT 1968
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Methods and materials are provided for the production of essentially
AΒ
       isomerically pure perhalogenated and partially halogenated compounds.
       One embodiment of the present invention provides a process for the
       production of essentially isomerically pure CFC-216aa. Other embodiments
       include processes for the production of CFC-217ba and HFC-227ea.
       Particular embodiments of the present invention provide separation
       techniques for the separation of chlorofluorocarbons from HF, from other
       chlorofluorocarbons, and the separation of isomers of halogenated
       compounds. Still other embodiments of the present invention provide
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catalytic synthetic techniques that demonstrate extended catalyst

techniques for the purification of isomeric mixtures.

lifetime. In other embodiments, the present invention provides catalytic

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L18 ANSWER 7 OF 12 USPATFULL on STN AN2004:9668 USPATFULL Processes for the purification and use of 2-chloro-1,1,1,2,3,3,3-TIheptafluoropropane and zeotropes thereof with HF INMiller, Ralph Newton, Newark, DE, United States Rao, V. N. Mallikarjuna, Wilmington, DE, United States Swearingen, Steven H., Wilmington, DE, United States E. I. du Pont de Nemours and Company, Wilmington, DE, United States PA (U.S. corporation) 20040113 See the See the US-6677493 B1 PIUS 1999-283449 AΙ US 1998-80709P 19980403 (60) PRAI DTUtility GRANTED FS Primary Examiner: Richter, Johann; Assistant Examiner: Price, Elvis O. EXNAM CLMN Number of Claims: 19 ECL Exemplary Claim: 1 DRWN 1 Drawing Figure(s); 1 Drawing Page(s) LN.CNT 610 CAS INDEXING IS AVAILABLE FOR THIS PATENT. A process is disclosed for the separation of a mixture of HF and AB

CF.sub.3CClFCF.sub.3. The process involves placing the mixture in a separation zone at a temperature of from about -30° C. to about 100° C. and at a pressure sufficient to maintain the mixture in the liquid phase, whereby an organic-enriched phase comprising less than 50 mole percent HF is formed as the bottom layer and an HF-enriched phase comprising more than 90 mole percent HF is formed as the top layer. The organic-enriched phase can be withdrawn from the bottom of the separation zone and subjected to distillation in a distillation column to recover essentially pure CF.sub.3CClFCF.sub.3. The distillate comprising HF and CF.sub.3CClFCF.sub.3 can be removed from the top of the distillation column while essentially pure CF.sub.3CClFCF.sub.3 can be recovered from the bottom of the distillation column. The HF-enriched phase can be withdrawn from the top of the separation zone and subjected to distillation in a distillation column. The distillate comprising HF and CF.sub.3CClFCF.sub.3 can be removed from the top of the distillation column while essentially pure HF can be recovered from the bottom of the distillation column. If desired, the two distillates can be recycled to the separation zone.

Also disclosed are compositions of hydrogen fluoride in combination with an effective amount of CF.sub.3CClFCF.sub.3 to form an azeotrope or azeotrope-like composition with hydrogen fluoride. Included are compositions containing from about 38.4 to 47.9 mole percent CF.sub.3CClFCF.sub.3.

Also disclosed are processes for producing 1,1,1,2,3,3,3-heptafluoro-propane. One process uses a mixture comprising HF and CF.sub.3CClFCF.sub.3 and is characterized by preparing essentially pure CF.sub.3CClFCF.sub.3 as indicated above, and reacting the CF.sub.3CClFCF.sub.3 with hydrogen. Another process uses an azeotropic composition as described above, and reacts the CF.sub.3CClFCF.sub.3 with hydrogen in the presence of HF.

Also disclosed is a process for producing hexafluoropropene. This process is characterized by preparing essentially pure CF.sub.3CClFCF.sub.3 as indicated above, and dehalogenating the CF.sub.3CClFCF.sub.3.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L18 ANSWER 8 OF 12 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 2 AN 2003:282507 CAPLUS

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Materials and methods for the production and purification of
ΤI
     chlorofluorocarbons and hydrofluorocarbons
     Iikubo, Yuichi; Owens, Stephen; Cohn, Mitchel; Brandstadter, Stephan M.;
IN
     Hedrick, Vicki E.; Boggs, Janet K.; Chien, John Chengping; Sacarias, Julie
     Pcbu Services, Inc., USA
PΑ
SO
     PCT Int. Appl., 66 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
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     PATENT NO.
                         KIND
                                                                     DATE
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     WO 2003029173
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     EP 1430010
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             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, SK
                                             U$ 2003-698579
     US 2004102661
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                                                                    20031031
                                             ŲS 2003-698730
     US 2004102662
                          Α1
                                20040527
                                                                    20031031
                                20040527
                                             ÚS 2003-698731
     US 2004102660
                          Α1
                                                                    20031031
                          Α1
                                            (US 2003-698923
     US 2004102663
                                20040527
                                                                    20031031
                                             US 2003-699491
     US 2004102664
                          A1
                                20040527
                                                                    20031031
                                20010928
PRAI US 2001-966158
                          Α
                          W
                                20020927 <
     WO 2002-US30729
OS
     CASREACT 138:289365
     Methods and materials are provided for the production of essentially
     isomerically pure perhalogenated and partially halogenated compds. One
     embodiment of the present invention provides a process for the production of
     essentially isomerically pure CFC-216aa. Other embodiments include
     processes for the production of CFC-217ba and HFC-227ea. Particular
     embodiments of the present invention provide separation techniques for the
     separation of chlorofluorocarbons from HF, from other chlorofluorocarbons, and
     the separation of isomers of halogenated compds. Still other embodiments of
     the present invention provide catalytic synthetic techniques that
     demonstrate extended catalyst lifetime. In other embodiments, the present
     invention provides catalytic techniques for the purification of isomeric mixts.
L18
     ANSWER 9 OF 12 USPATFULL on STN
AN
       2003:153700 USPATFULL
TI
       Materials and methods for the production and purification of
       chlorofluorocarbons and hydrofluorocarbons
IN
       Iikubo, Yuichi, West Lafayette, IN, UNITED STATES
       Owens, Stephen, White Pine, TN, UNITED STATES
       Cohn, Mitchel, West Lafayette, IN, UNITED STATES
       Brandstadter, Stephan M., Indianapolis, IN, UNITED STATES
       Hedrick, Vicki E., Brookston, IN, UNITED STATES
       Boggs, Janet K., Brownsburg, IN, UNITED STATES
       Qian, John, West Lafayette, IN, UNITED STATES
```

Sacarias, Julie, El Dorado, AR, UNITED STATES

20030605

20010928 (9)

Α1

A1

138:289365

DN

PI

ΑI

US 2003105368

US 2001-966158

```
Utility
DT
FS
       APPLICATION
       BAKER & DANIELS, 300 NORTH MERIDIAN STREET, SUITE 2700, INDIANAPOLIS,
LREP
       IN, 46204-1782
       Number of Claims: 88
CLMN
       Exemplary Claim: 1
ECL
       7 Drawing Page(s)
DRWN
LN.CNT 2001
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Methods and materials are provided for the production of essentially
AB
       isomerically pure perhalogenated and partially halogenated compounds.
       One embodiment of the present invention provides a process for the
       production of essentially isomerically pure CFC-216aa. Other embodiments
       include processes for the production of CFC-217ba and HFC-227ea.
       Particular embodiments of the present invention provide separation
       techniques for the separation of chlorofluorocarbons from HF, from other
       chlorofluorocarbons, and the separation of isomers of halogenated
       compounds. Still other embodiments of the present invention provide
       catalytic synthetic techniques that demonstrate extended catalyst
       lifetime. In other embodiments, the present invention provides catalytic
       techniques for the purification of isomeric mixtures.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L18 ANSWER 10 OF 12 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 3
     2001:490662 CAPLUS
AN
DN
     135:241863
     13C NMR spectra of halocarbons
TI
ΑU
     Foris, Anthony
     Jackson Laboratory, DuPont Central Research and Development, Deepwater,
CS
     NJ, 08203, USA
     Magnetic Resonance in Chemistry (2001), 39(7), 386-398
SO
     CODEN: MRCHEG; ISSN: 0749-1581
     John Wiley & Sons Ltd.
PB
DT
     Journal
LA
     English
     13C NMR chemical shifts and 13C-19F and 13C-1H coupling consts. are reported
AB
     for 103 halocarbons. Methods used for analyzing complex spectra (e.g.
     simultaneous 1H and 19F broadband decoupling, 2-dimensional
     INAD-EQUATE/FRED, HMQC, 35,37Cl isotope effects) are briefly illustrated.
     Longitudinal relaxation times for CF3CCl2CH2CHClF are reported.
              THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 46
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
L18 ANSWER 11 OF 12 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 4
     1999:659342 CAPLUS
AN
DN
     131:272325
     Processes for the distillative purification and use of
TI
     2-chloro-1,1,1,2,3,3,3-heptafluoropropane and its azeotropes with HF in
     the manufacture of hexafluoropropene and 1,1,1,2,3,3,3-heptafluoropropane
     Miller, Ralph Newton; Rao, V. N. Mallikarjuna; Swearingen, Steven H.
IN
     E. I. Du Pont de Nemours & Co., USA
PA
     PCT Int. Appl., 18 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
                                                                    DATE
                                            APPLICATION NO.
    PATENT NO
                         KIND
                                DATE
                                                                    19990401
                                19991014
                                            WO 1999-US7225
     WO 9951555
PI
                          A1
         W:-AE, AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GD, GE, HR, HU,
             ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MG, MK, MN, MX,
             NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
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RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, .
            ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
            CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                       AU 1999-33779
                                                                19990401
                              19991025
    AU 9933779
                        A1
                                                                19990401
                                         EP 1999-915210
                              20010110
                        A1
    EP 1066230
                        B1
                              20030910
    EP 1066230
        R: BE, DE, ES, FR, GB, IT, NL
                                         JP 2000-542278
                                                                19990401
                       T2
                              20020409
    JP 2002510664
                                                                19990401
                                       US 1999-283449
                              20040113
                        B1
   US 6677493
                                         ES 1999-915210
                                                                19990401
                        T3
                              20040401
    ES 2203107
PRAI US 1998-80709P
                       P
                              19980403
                              19990401
                        W
    WO 1999-US7225
```

The separation of a mixture of HF and CF3CClFCF3 involves placing the mixture AΒ in a

separation zone at a temperature of from about -30° to about +100° and at a pressure sufficient to maintain the mixture in the liquid phase, so that an organic-enriched phase comprising <50 mol percent HF is formed as the bottom layer and an HF-enriched phase comprising >90 mol percent HF is formed as the top layer. The organic-enriched phase is withdrawn from the bottom of the separation zone and subjected to distillation in a distillation column to

recover essentially pure CF3CClFCF3. The distillate comprising HF and CF3CClFCF3 can be removed from the top of the distillation column while essentially pure CF3CClFCF3 can be recovered from the bottom of the distillation

The HF-enriched phase can be withdrawn from the top of the separation column. zone and subjected to distillation in a distillation column. The distillate comprising HF and CF3CC1FCF3 can be removed from the top of the distillation column while essentially pure HF can be recovered from the bottom of the distillation column. If desired, the two distillates can be recycled back to the separation zone. Also disclosed are compns. of hydrogen fluoride in combination with an effective amount of CF3CClFCF3 to form an azeotrope-like composition with HF; included are compns. containing 38.4-47.9 mol percent CF3CClFCF3. Also disclosed are processes for producing

1,1,1,2,3,3,3-heptafluoropropane and hexafluoropropene.

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 5 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L18 ANSWER 12 OF 12 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 5

1975:427097 CAPLUS AΝ

DN83:27097

Mass spectral correlations of halogenated 1,1,1,3,3,3-hexafluoropropanes ŢΙ and 1,1,1-trifluoroethanes

Naae, Douglas G.; Wiebe, Donald A. ΑU

Dep. Chem., Univ. Iowa, Iowa City, IA, USA CS

Organic Mass Spectrometry (1974), 9(12), 1203-6 SO CODEN: ORMSBG; ISSN: 0030-493X

DTJournal

English LΑ

The mass spectra of 12 CF3CXYCF3 (X and Y = H, F, Cl, Br, and I) and 5 AB CF3CX2Y (X and Y = F, Cl, and Br) were recorded at 20 and 70 eV and the observed trends were discussed. The frequency of carbon-halogen and C-C bond cleavage was dependent on the substituents X and Y.

```
ANSWER 1 OF 4 USPATFULL on STN
L23
       97:31869 USPATFULL
AN
       Halocarbon hydrogenolysis
TI
       Manogue, William H., Newark, DE, United States
IN
      E. I. Du Pont De Nemours and Company, Wilmington, DE, United States
PA
                            2 19970415
       (U.S. corporation)
      ÚS 562115]
PI
                               19950421 (8)
       US 1995-426534
ΑI
       Division of Ser. No. US 1994-242503, filed on 13 May 1994, now patented,
RLI
       Pat. No. US 5430204 which is a continuation of Ser. No. US 1993-122102,
       filed on 16 Sep 1993, now patented, Pat. No. US 5364992 And a
       continuation of Ser. No. US 1992-847987, filed on 9 Apr 1992, now
       abandoned
       Utility
\mathbf{DT}
FS
       Granted
EXNAM Primary Examiner: Siegel, Alan
       Number of Claims: 20
CLMN
       Exemplary Claim: 1
ECL
DRWN
       No Drawings
LN.CNT 791
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Halocarbons such as CCl.sub.2 F.sub.2, CF.sub.3 CFHCl or CF.sub.3
       CFCl.sub.2 which contain chlorine and/or bromine are contacted with
       hydrogen in the presence of silicon carbide and/or a metal selected from
       aluminum, molybdenum, titanium, nickel, iron or cobalt (or their alloys)
       at temperatures of 350° to 700° C. and pressures of 0 to
       1000 psig to obtain a product wherein at least one chlorine or bromine
       in the starting material has been replaced by hydrogen.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 2 OF 4 USPATFULL on STN
L23
       95:60539 USPATFULL
AN
       Halocarbon hydrogenolysis
TI
       Manogue, William H., Newark, DE, United States
IN
       Rao, V. N. Mallikarjuna, Wilmington, DE, United States
       E. I. Du Pont de Nemours and Company, Wilmington, DE, United States
PΑ
       (O.S. corporation)
                               19950704
19940513 (8) See Als
       US-543.0204
PI
       US 1994-242503
AΙ
       20111115
DCD
       Continuation of Ser. No. US 1993-122102, filed on 16 Sep 1993, now
RLI
       patented, Pat. No. US 5364992 which is a continuation of Ser. No. US
       1992-847987, filed on 9 Apr 1992, now abandoned which is a
       continuation-in-part of Ser. No. US 1989-418832, filed on 10 Oct 1989,
       now abandoned
DT
       Utility
       Granted
FS
       Primary Examiner: Siegel, Alan
EXNAM
       Number of Claims: 20
CLMN
       Exemplary Claim: 1
ECL
       No Drawings
DRWN
LN.CNT 829
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Halocarbons such as CCl.sub.2 F.sub.2, CF.sub.3 CFHCl or CF.sub.3
AB
       CFCl.sub.2 which contain chlorine and/or bromine are contacted with
       hydrogen in the presence of silicon carbide and/or a metal selected from
       aluminum, molybdenum, titanium, nickel, iron or cobalt (or their alloys)
       at temperatures of 350° to 700° C. and pressures of 0 to
       1000 psig to obtain a product wherein at least one chlorine or bromine
       in the starting material has been replaced by hydrogen.
```

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94:100045 USPATFULL
AN
      Halocarbon hydrogenolysis
TI
      Manoque, William H., Newark, DE, United States
IN
       Rao, Velliyur N. M., Wilmington, DE, United States
       E. I. Du Pont de Nemours and Company, Wilmington, DE, United States
PA
       (U.S. corporation)
                              19941115 Lee this
19930916 (8)
       US 5364992 /
PI
      US 1993-122102
ΑI
      Continuation of Ser. No. US 1992-847987, filed on 9 Apr 1992, now
RLI
       abandoned which is a continuation-in-part of Ser. No. US 1989-418832,
       filed on 10 Oct 1989, now abandoned
      Utility
DT
      Granted
FS
      Primary Examiner: Siegel, Alan
EXNAM
      Number of Claims: 18
CLMN
ECL
      Exemplary Claim: 1
      No Drawings
DRWN
LN.CNT 789
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Halocarbons such as CCl.sub.2 F, CF.sub.3 CFHCl.sub.2 or CF.sub.3
AB
       CFCl.sub.2 which contain chlorine and/or bromine are contacted with
      hydrogen in the presence of silicon carbide and/or a metal selected from
       aluminum, molybdenum, titanium, nickel, iron or cobalt (or their alloys)
       at temperatures of 350° to 700° C. and pressures of 0 to
       1000 psig to obtain a product wherein at least one chlorine or bromine
       in the starting material has been replaced by hydrogen.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L23 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 1
     1991:558497 CAPLUS
AN
     115:158497
DN
     Halocarbon hydrogenolysis
TI
     Kielhorn, Fernando Frederick; Manogue, William Henry; Rao, V. N.
IN
     Mallikarjuna
     du Pont de Nemours, E. I., and Co., USA
PA
     PCT Int. Appl., 33 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
                               DATE
                                          APPLICATION NO.
                                                                  DATE
     PATENT NO.
                        KIND
     ______
                         - - - -
                                           WO 1990-US5637
                               19910502
                         A1
                                                                  19901009
     WO 9105752
PI
        W: AU, BR, CA, JP, KR, SU, US
        RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE
                               19950603
                                         IN 1990-CA857
                                                                  19901008
     IN 175284
                         Α
                                           CA 1990-2067370
                                                                  19901009
     CA 2067370
                        AA
                               19910411
                                         AU 1990-66113
                                                                  19901009
     AU 9066113
                         A1
                               19910516
     AU 641704
                        B2
                               19930930
                                         ZA 1990-8087
                                                                  19901009
                        Α
                               19920624
     ZA 9008087
                                         EP 1990-915920
                                                                  19901009
                         A1
                               19920729
     EP 495892
     EP 495892
                        В1
                               19950719
        R: DE, ES, GB, IT, NL
                                                                  19901009
                                           BR 1990-7726
     BR 9007726
                         Α
                               19920915
                                           JP 1990-514986
                                                                  19901009
                         T2
                               19930408
     JP 05501878
     JP 2825650
                         B2
                               19981118
                                19951001
                                           ES 1990-915920
                                                                  19901009
     ES 2075225
                         Т3
                                           RU 1990-5011233
                         C1
                                19961110
                                                                  19901009
     RU 2068835
                                           CN 1990-109466
                                                                  19901010
     CN 1051167
                                19910508
                         Α
     CN 1029119
                         В
                               19950628
                                           US 1993-122102
     US 5364992
                                19941115
                                                                  19930916
                         Α
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ANSWER 3 OF 4 USPATFULL on STN

L23

	US 5430204	A	19950704	US	1994-242503	19940513
	US 5621151	A	19970415	US	1995-426534	19950421
PRA:	I US 1989-418832	A2	19891010			
	WO 1990-US5637	A	19901009			
	US 1992-847987	B1	19920409			tU
	US 1993-122102	A1	19930916			
	US_1994-242503	A3	19940513) le	
OS	CASREACT 115:158497:	MARPA	Т 115:158497		0	

A process for the hydrogenolysis of halocarbons of the general formula CnHmFpXq wherein n = 1-10, m = 0-20, p = 0-21, q = 1-22, X = Cl, Br comprises reacting said halocarbon with hydrogen in a vessel packed with particles of Al, Mo, Ti, Ni, Fe, Co or their alloys or silicon carbide or low surface area carbon at high pressures and temps. Thus, CF3CCl2F (1.47 g/h) and hydrogen (molar ratio of H2/CF3CCl2F = 1.9) were fed into a 1/4" empty Inconel Ni alloy reactor for 38 h at 450-550° and 250 psig. A sample taken after 14 h at 550° showed 89% conversion of CF3CCl2F with 65% selectivity to CF3CHClF and 32% selectivity to CF3CH2F. Overall selectivity of 2 products was 97%.

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ANSWER 1 OF 5 USPATFULL on STN
L25
       2002:6200 USPATFULL
\mathbf{A}\mathbf{N}
       Circuit arrangement and method for operating and electrical motor
\mathtt{TI}
       Gleim, Gunter, Villingen-Schwenningen, GERMANY, FEDERAL REPUBLIC OF
IN
       Deutsche Thomson-Brandt GmbH, Villengen-Schwenningen, GERMANY, FEDERAL
PA
       REPUBLIC OF (non-U.S. corporation)
                          B1
                               20020108
PI
       US 6337553
                               19980518 (9)
       US 1998-80708
AI
                        19970523
      DE 1997-19721490
PRAI
      Utility
DT
FS
       GRANTED
      Primary Examiner: Masih, Karen
EXNAM
       Tripoli, J. S., Herrmann, E. P.
LREP
CLMN
       Number of Claims: 6
       Exemplary Claim: 1
\mathsf{ECL}
       3 Drawing Figure(s); 1 Drawing Page(s)
DRWN
LN.CNT 165
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       A circuit arrangement with a digital controller for operating an
       electric motor and a method for starting an electric motor are
       disclosed. Electric motors require a considerably higher current
       immediately after they have been switched on and during the running-up
       phase because of the inductance of the motor windings. In order to
       increase the control sensitivity of the controller for the normal
       operation, the control range of a digital controller is switched over
       into a higher current range for the running-up phase. For the normal
       operation of the motor, the control range of the controller is switched
       back again into a lower current range. As a result, the control
       sensitivity of the controller is completely available for the normal
       operation of the motor.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
    ANSWER 2 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 1
L25
     2001:630912 CAPLUS
AN
     135:182382
DN
     1,1,1,2,3,3,3-Heptafluoropropane manufacturing process
TI
    Nappa, Mario Joseph; Rao, V. N. Mallikarjuna; Sievert, Allen Capron
IN
     E. I. Du Pont de Nemours & Co., USA
PΑ
     U.S., 5 pp.
SO
     CODEN: USXXAM
DT
     Patent
LA
     English
FAN.CNT 1
                                DATE
                                            APPLICATION NO.
                                                                    DATE
     PATENT NO.
                                            _______
                                                                    _ _ _ _ _ _ _ _ _ _
     ____
                                            US 1999-283451
                                                                  19990401
                          В1
                                20010828
    US 6281395/
PI'
                          P
                                19980403
PRAI US-1998-80706P
     A process is disclosed for the manufacture of CF3CHFCF3 containing <0.01 ppm
AΒ
     (CF3)2C:CF2. The process involves: (a) contacting hexafluoropropene in
     the vapor phase at <260° with hydrogen fluoride in the presence of
     a selected fluorination catalyst or produce a product containing <10 parts
     (CF3)2C:CF2 per million parts of CF3CHFCF3; and (b) treating the product
     of (a) as necessary to remove excess (CF3)2C:CF2. Suitable catalysts
     include: (i) an activated carbon treated to contain from about 0.1-10% of
     added alkali or alkaline earth metals; (ii) three dimensional matrix porous
     carbonaceous materials; (iii) supported metal catalysts comprising
     trivalent chromium; and (iv) unsupported chrome oxide prepared by the
     pyrolysis of (NH4)2Cr2O7.
```

THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

AN 2000:10072 USPATFULL

TI Process for the production of fluorocarbons

IN Manogue, William H., Newark, DE, United States Nappa, Mario Joseph, Newark, DE, United States Sievert, Allen Capron, Elkton, MA, United States

Rao, V. N. Mallikarjuna, Newark, DE, United States

PA E. I. du Pont de Nemours and Company, Wilmington, DE, United States

Sa Huz

(U.S. corporation)

US 1999-283450 20000125 19990401 (9)

AI US 1999-283450 PRAI US 1998-80708P

19980403 (60)

DT Utility

OCTITCY

FS Granted

EXNAM Primary Examiner: Siegel, Alan

CLMN Number of Claims: 3

ECL Exemplary Claim: 1
DRWN 1 Drawing Figure(s); 1 Drawing Page(s)

be recycled to the separation zone.

LN.CNT 495

PΙ

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A process is disclosed for the separation of a mixture of HF and AB CF.sub.3 CClFCF.sub.3. The process involves placing the mixture in a separation zone at a temperature of from about -30° C. to about 100° C. and at a pressure sufficient to maintain the mixture in the liquid phase, whereby an organic-enriched phase comprising less than 50 mole percent HF is formed as the bottom layer and an HF-enriched phase comprising more than 90 mole percent HF is formed as the top layer. The organic-enriched phase can be withdrawn from the bottom of the separation zone and subjected to distillation in a distillation column to recover essentially pure CF.sub.3 CClFCF.sub.3. The distillate comprising HF and CF.sub.3 CClFCF.sub.3 can be removed from the top of the distillation column while essentially pure CF.sub.3 CClFCF.sub.3 can be recovered from the bottom of the distillation column. The HF-enriched phase can be withdrawn from the top of the separation zone and subjected to distillation in a distillation column. The distillate comprising HF and CF.sub.3 CClFCF.sub.3 can be removed from the top of the distillation column while essentially pure HF can be recovered from the bottom of the distillation column. If desired, the two distillates can

Also disclosed are compositions of hydrogen fluoride in combination with an effective amount of CF.sub.3 CClFCF.sub.3 to form an azeotrope or azeotrope-like composition with hydrogen fluoride. Included are compositions containing from about 38.4 to 47.9 mole percent CF.sub.3 CClFCF.sub.3.

Also disclosed are processes for producing 1,1,1,2,3,3,3-heptafluoropropane. One process uses a mixture comprising HF and CF.sub.3 CCIFCF.sub.3 and is characterized by preparing essentially pure CF.sub.3 CCIFCF.sub.3 as indicated above, and reacting the CF.sub.3 CCIFCF.sub.3 with hydrogen. Another process uses an azeotropic composition as described above, and reacts the CF.sub.3 CCIFCF.sub.3 with hydrogen in the presence of HF.

Also disclosed is a process for producing hexafluoropropene. This process is characterized by preparing essentially pure CF.sub.3 CClFCF.sub.3 as indicated above, and dehalogenating the CF.sub.3 CClFCF.sub.3.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L25 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 2

AN 1999:659340 CAPLUS

DN 131:258061

TI Process for the production of hexafluoropropylene and 1,1,1,2,3,3,3-

```
Manogue, William H.; Nappa, Mario Joseph; Sievert, Allen Capron
IN
     E. I. Du Pont de Nemours & Co., USA
PA
SO
     PCT Int. Appl., 16 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
                                            APPLICATION NO.
                                                                   DATE
                                DATE
     PATENT-NO.
                                            _____.
                                                                   19990401
                                            WO 1999-US7230
     WO 9951553
                          A1
                                19991014
PΙ
             AE, AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GD, GE, HR, HU,
        W:
             ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MG, MK, MN, MX,
             NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                19991025
                                          AU 1999-33780
                                                                   19990401
     AU <u>99</u>33780
                          A1
                                            US 1999-283450
                                                                   19990401
    US 6018083
                          Α
                                20000125
                                            EP 1999-915212
                                                                   19990401
     EP 1068167
                                20010117
                          A1
                          В1
                                20030903
     EP 1068167
         R: BE, DE, ES, FR, GB, IT, NL
                                                                   19990401
                         T2
                                            JP 2000-542276
     JP 2002510662
                                20020409
                                            ES 1999-915212
                                                                   19990401
                          Т3
                                20040401
     ES 2203108
                          Ρ
                                19980403
PRAI US 1998-80708P
                          W
                                19990401
     WO 1999-US7230
     Hexafluoropropylene and 1,1,1,2,3,3,3-heptafluoropropane are manufactured by:
AΒ
     (A) feeding 1,1,2-trichloro-3,3,3-trifluoro-1-propene, HF, and Cl2 to a
     first reaction zone containing a trivalent chromium catalyst operated at
     250-325° to produce an effluent comprising C3Cl3F5, C3Cl2F6,
     CF3CClFCF3, HCl, and HF; (B) the effluent of step A is distilled to produce
     (i) a low-boiling stream including HCl, (ii) a reactant stream including
     an azeotrope of 2-chloro-1,1,1,2,3,3,3-heptafluoropropane and HF, and
     (iii) a high-boiling stream including C3Cl2F6 and C3Cl3F5; (C)
     2-chloro-1,1,1,2,3,3,3-heptafluoropropane of reactant stream (ii) is
     reacted with hydrogen in the presence of a catalyst to produce a mixture of
     hexafluoropropylene and 1,1,1,2,3,3,3-heptafluoropropane; (D) the C3Cl2F6
     and C3Cl3F5 of high-boiling stream (iii) are fed along with HF to a second
     reaction zone containing a trivalent chromium catalyst and operated at
     ≥375° to produce a reaction product comprising CF3CClFCF3
     and HF; and (E) the product mixture of step D is recycled to step A. A
     process flow diagram is presented.
RE.CNT 2
              THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 5 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 3
L25
AN
     1993:59239 CAPLUS
DN
     118:59239
     Reaction of organic compounds with a sulfur tetrafluoride-hydrogen
TI
     fluoride-halogenating agent system. VII. Reactions of olefins with the
     SF4-HF-Cl2(Br2) system
     Kunshenko, V. B.; Mohamed, Nagib Muhtar; Omarov, V. O.; Muratov, N. N.;
ΑU
     Yagupol'skii, L. N.
     Odess. Politekh. Inst., Odessa, Ukraine
CS
     Zhurnal Organicheskoi Khimii (1992), 28(4), 672-80
SO
     CODEN: ZORKAE; ISSN: 0514-7492
DT
     Journal
     Russian
LA
     CASREACT 118:59239
OS
     Halogenated alkenes undergo halofluorination in SF4-HF-Cl2(Br2) systems.
AB
     On the basis of Z- and E-1,2-dichloroethenes it was shown that these
     reactions proceed with anti stereospecificity via bromonium ions. The
     accumulation of Cl atoms in the alkene mol. hinders electrophilic addition of
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heptafluoropropane

stoichiometric equivs. of ClF and BrF to the double bond. The SF4-HF-Br2 system is effective in fluorinating Br-containing organic compds., wherein only Br atoms on a secondary C are substituted by F.

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ANSWER 1 OF 2 CAPLUS COPYRIGHT 2004 ACS on STN
L32
     2004:534048 CAPLUS
AN
     141:89877
DN
     Materials and methods for the conversion of hydrofluorocarbons to
TI
     fluoromonomers
     Iikubo, Yuichi; Hedrick, Vicki; Brandstadter, Stephen M.; Cohn, Mitchel
IN
PA
     U.S. Pat. Appl. Publ., 11 pp.
SO
     CODEN: USXXCO
     Patent
DT
     English
LA
FAN.CNT 1
                                                                    DATE
                                            APPLICATION NO.
                         KIND
                                DATE
     PATENT NO.
                         _ _ _ _
                                20040701
                                                                    20021230
                                            US 2002-331821
                          A1
PΙ
     US 2004127757
                                                                    20031230
                                20040722
                                            WO 2003-US41851
                          A1
     WO 2004060842
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
         W:
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
             PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
             TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU
         RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE,
             BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU,
             MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN,
             GQ, GW, ML, MR, NE, SN, TD, TG
PRAI US 2002-331821
                          A2
                                20021230
     Methods and materials for the recovery of valuable hydrofluorocarbons and
AB
     subsequent conversion to fluoromonomer precursers and fluoromonomers are
     disclosed. More specifically methods and materials are provided for
     recovering hydrofluorocarbons such as HFC-227, HFC-236, HFC-245, HFC-125,
     HFC-134, HFC-143, HFC-152, HFC-32, HFC-23 and their resp. isomers.
     Processes are provided for converting hydrofluorocarbons such as these to
     fluoromonomer precursors such as CFC-217, CFC-216, CFC-215, CFC-115,
     CFC-114, CFC-113, CFC-112, HCFC-22, CFC-12, CFC-13 and their resp.
     isomers. Materials, methods, and schemes are provided for the conversion
     of these fluoromonomer precursors to fluoromonomers such as HFP, PFP, TFP,
     TFE, and VDF. One example demonstrates the conversion of HFC-227 to
     CFC-217 and finally to hexafluoropropene.
     ANSWER 2 OF 2 CAPLUS COPYRIGHT 2004 ACS on STN
L32
     1986:552494 CAPLUS
AN
     105:152494
DN
     Interaction of fluoro olefins with halogens in highly acid medium
TI
     Fokin, A. V.; Studnev, Yu. N.; Rapkin, A. I.; Krotovich, I. N.; Tatarinov,
ΑŲ
     A. S.; Verenikin, O. V.
     Inst. Elementoorg. Soedin., Moscow, USSR
CS
     Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1985), (10), 2298-302
SO
     CODEN: IASKA6; ISSN: 0002-3353
DT
     Journal
     Russian
LA
os
     CASREACT 105:152494
     Halogenation of F2C:CFR (R = F, CF3) in HSO3F or HSO3F/SbF5 gave
AB
     RCFXCF2OSO2F (X = Cl, Br, iodo), RCFXCF2X (X = Cl, Br), RCFICF3, (CF3)2CFX
     (X = Cl, Br) and F3CCHFCF2OSO2F in yields depending on reaction
     conditions.
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ANSWER 1 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
    2002:539627 CAPLUS
\mathbf{A}\mathbf{N}
DN
    137:95530
    Process for manufacturing and purifying octafluorocyclobutane
TI
    Horiba, Minako; Suzuki, Yasuhiro
IN
PA
    Showa Denko K.K., Japan
SO
    PCT Int. Appl., 56 pp.
    CODEN: PIXXD2
    Patent
DT
    English
LA
FAN.CNT 1
                       KIND DATE
                                         APPLICATION NO.
                                                                  DATE
    PATENT NO.
                                           _____
                        - - - -
                               _____
                                                                  20020111
                        A2
                                           WO 2002-JP148
PΙ
    WO 2002055458
                               20020718
                        A3
                               20030220
    WO 2002055458
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
            LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL,
            PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA,
            UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                        JP 2001-6459
                                                                 20010115
    JP 2002212118
                         A2
                               20020731
                               20030507
                                           EP 2002-729558
                                                                  20020111
                         A2
    EP 1307416
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                          US 2002-221443
                                                                  20020912
                        A1
                               20030717
    US 2003132099
                         Α
                               20010115
PRAI JP 2001-6459
                         Ρ
                               20010129
    US 2001-264322P
                         W
                               20020111
    WO 2002-JP148
    Octafluorocyclobutane is purified by contacting crude
AΒ
    octafluorocyclobutane containing impurities with an impurity decomposing agent
at
    elevated temperature and then with an adsorbent to substantially remove the
     impurities from the crude octafluorocyclobutane. The obtained
    octafluorocyclobutane is substantially free of impurities and therefore,
     can be used as an etching or cleaning gas in the production of semiconductor
     devices or the like.
    ANSWER 2 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
L33
AN
    1999:42330 CAPLUS
DN
    130:112046
     Gas chromatographic retention parameters database for refrigerant mixture
TI
     composition management
    Bruno, Thomas J.; Bachmeyer, Gregory M.; Wertz, Kelly H.
ΑU
     Physical and Chemical Properties Division, Chemical Science and Technology
CS
    Laboratory, National Institute of Standards and Technology, Boulder, CO,
     80303, USA
     International Journal of Refrigeration (1998), 21(8), 639-647
SO
     CODEN: IJRFDI; ISSN: 0140-7007
    Elsevier Science Ltd.
PB
DT
    Journal
LA
     English
     Composition management of mixed refrigerant systems is a challenging problem in
AB
     the laboratory, manufacturing facilities, and large refrigeration machinery.
     issue of composition management is especially critical for the maintenance of
machinery
     that utilizes zeotropic mixts. as working fluids. These are fluids in
    which the gas and liquid phases will generally have greatly different
     compns. While there are many anal. techniques available for laboratory and
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online analyses, gas chromatog. probably offers the greatest flexibility at the most reasonable cost. This paper describes a chromatog. database that provides for the identification of refrigerant components, and thereby facilitates composition management of zeotropic fluids. Prior to the description of the database a description is given of the basic theory of chromatog. retention parameters and the exptl. techniques used in their measurement.

- RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L33 ANSWER 3 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1996:148274 CAPLUS
- DN 124:249231
- TI Kovats Retention Indexes of Halocarbons on a Hexafluoropropylene Epoxide-Modified Graphitized Carbon Black
- AU Bruno, Thomas J.; Wertz, Kelly H.; Caciari, Michael
- CS Thermophysics Division, National Institute of Standards and Technology, Boulder, CO, 80303, USA
- SO Analytical Chemistry (1996), 68(8), 1347-59 CODEN: ANCHAM; ISSN: 0003-2700
- PB American Chemical Society
- DT Journal
- LA English
- Kovats retention indexes of 97 halocarbons related to research on AB alternative refrigerants, propellants, foaming agents, and blowing agents were measured on a packed column stationary phase consisting of a 5% (mass/mass) coating of a low mol. weight polymer of hexafluoropropylene epoxide on graphitized carbon black. The measurements on each fluid were made at four temps., and the thermal dependence of the indexes was modeled with appropriate equations. The modeled values are suitable for the identification of these compds. by gas chromatog. on both laboratory and field instrumentation. The values are also useful for the optimization of more sophisticated analyses needed in specific situations. The stationary phase chosen will provide separation of nearly all the fluids of interest. Also, there is sufficient spread in the retention index values to facilitate fluid identification. The measurements also appear to fit a qual. triangular property diagram that was useful for classifying alternative refrigerant fluids and related compds.
- L33 ANSWER 4 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1995:739446 CAPLUS
- DN 123:305689
- TI Retention of halocarbons on a hexafluoropropylene epoxide-modified graphitized carbon black. IV. Propane-based compounds
- AU Bruno, Thomas J.; Wertz, Kelly H.; Caciari, Michael
- CS Thermophysics Division, Chemical Science and Technology Laboratory, National Institute of Standards and Technology, Boulder, CO, 80303, USA
- SO Journal of Chromatography, A (1995), 708(2), 293-302 CODEN: JCRAEY; ISSN: 0021-9673
- PB Elsevier
- DT Journal
- LA English
- AB The retention characteristics of 25 propane-based bromofluorocarbon, chlorocarbon, chlorofluorocarbon, and fluorocarbon fluids were studied as a function of temperature on a stationary phase consisting of a 5% (m/m)
 - of a low-mol.-mass polymer of hexafluoropropylene epoxide on a graphitized carbon black adsorbent. Measurements were performed at 0, 20, 40 and 60° for R-245ca and R-245cb. Measurements were performed at 20, 40, 60 and 80° for R-227ca, R-227ea, R-236ea, R-236fa, R-245fa, and R-263fb. Measurements were performed at 40, 60, 80 and 100° for R-217ba, R-254cb and R-1243b, and at 60, 80, 100 and 120° for R-280da and R-217caB1. Measurements were performed at 80, 100, 120 and 140° for R-215aa, R-216ba, R-253fb, R-262da, and R-270aa.

Measurements were performed at 100, 120,140 and 160° for R-215ba, R-225ca, R-225cb, R-243db, R-270da, R-270fa, and R-270fb. Relative retentions as a function of temperature were calculated with respect to the retentions of tetrafluoromethane (R-14) and hexafluoroethane (R-116). Qual. features of the data are examined, and trends are identified. The relative retention data were fitted to linear models for the purpose of predicting retention behavior of these compds. to facilitate chromatog. anal.

L33 ANSWER 5 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1993:516774 CAPLUS

DN 119:116774

TI Preparation of 1,1,1,2,3,3,3-heptafluoropropane (R 227) via catalytic hydrodechlorination of chloroheptafluoropropane

IN Hopp, Peter

PA Hoechst A.-G., Germany

SO Eur. Pat. Appl., 3 pp. CODEN: EPXXDW

DT Patent

LA German

FAN CNT 1

FAN.CNI I	Y			
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
(
PI EP 539989	A1	19930505	EP 1992-118515	19921029
EP 539989	B1	19970618		
R: BE, CH, DE,	DK, ES	, FR, GB, IT	, LI, NL, SE	
ES 210 <u>47</u> 90	Т3	19971016	ES 1992-118515	19921029
CA 2081813	ĀA	19930502	CA 1992-2081813	19921030
CA 20 81813	С	20040113		
JP 05221894	A2	19930831	JP 1992-293383	19921030
JP 3249202	B2	20020121		
PRAI DE 1991-4136054	A	19911101		

AB CF3CHFCCF3 was prepared by hydrodechlorination of CF3CClFCF3 (I) using the catalysts Pt, Pd, Ru, Rh, Ni, or Cu at 100-400° and 1-50 bar H2. The molar ratio of I with H2 was 1:1 to 1:10.

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L33 ANSWER 6 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
     1982:34523 CAPLUS
AN
DN
     96:34523
     2-Chloro-1,1,1,2,3,3,3-heptafluoropropane by photochlorination of
TI
     1,1,1,2,3,3,3-heptafluoropropane
     Von Halasz, Sigmar Peter
IN
PA
     Hoechst A.-G. , Fed. Rep. Ger.
     Ger. Offen., 14 pp.
SO
     CODEN: GWXXBX
DT
     Patent
     German
LA
FAN.CNT 1
                                DATE
                                            APPLICATION NO.
                                                                   DATE
     PATENT NO.
                         KIND
                         _ _ _ _
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                                _ _ _ _ _ _ _ _
                                19811112 DE 1980-3017154
PI
     DE 3017154
                          A1
                                                                    19800505
                                19820216 ES 1981-501729
     ES 501729
                         A1
                                                                   19810428
                                19811111 EP 1981-103200
     EP 39471
                          Al
                                                                    19810429
         R: BE, DE, FR, GB, IT, NL
     JP 56169632
                                            JP 1981-65307
                         A2
                                19811226
                                                                    19810501
                                            BR 1981-2724
     BR 8102724
                          Α
                                19820126
                                                                    19810504
                                            CA 1981-376831
                                19840221
                                                                    19810504
     CA 1162511
                          A1
PRAI DE 1980-3017154
                                19800505
     CF3CFClCF3 (I) was prepared by photochem. chlorination of CF3CHFCF3 (II) at
AB
     -30 to +500° (especially 40-450°) and a II-Cl Mol ratio 1:1.0-1.5,
     especially 1:1.05-1.15; a typical product contained 98.1% I.
L33 ANSWER 7 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
     1965:488299 CAPLUS
AN
     63:88299
DN
OREF 63:16189b-c
     Influence of substitution on CH.CF and CF.CF coupling constants
TI
     Abraham, R. J.; Cavalli, L.
ΑU
CS
     Univ. Liverpool, UK
     Molecular Physics (1965), 9(1), 67-75
SO
     CODEN: MOPHAM; ISSN: 0026-8976
     Journal
DT
     English
LA
AB
     The dependence of the coupling constant in CH.CF and CF.CF fragments on the
     electronegativities of the substituents, in the light of all acceptable
     data, was investigated, JHFAV obeyed the equation JFFAV = 53.03-3.38
     \Sigma E, while JHFAV \approx 91.4-6.15 \Sigma E, where \Sigma E is the
     sum of electronegativities of the 1st atom of the remaining substituents.
     Consideration of these equations, and of that for JHHAV showed that for
     highly electroneg. substituents all 3 couplings decrease to 0. The temperature
     dependence of the F-F couplings in some unsym. substituted ethanes was
     found due to the change in jFF with temperature and not the changing
populations
     of the rotational isomers.
                                 29 refs.
L33 ANSWER 8 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
AN
     1956:19880 CAPLUS
     50:19880
DN
OREF 50:4048d-h
     The reactions of highly fluorinated organic compounds. IX.
     1H-Decafluoro-4-trifluoromethylcyclohexane, nonafluoro-4-trifluoromethyl-1-
     cyclohexene, and perfluoro-(3-methyladipic) acid
ΑU
     Barlow, G. B.; Stacey, M.; Tatlow, J. C.
     Univ. Birmingham, UK
CS
     Journal of the Chemical Society, Abstracts (1955) 1749-52
SO
     CODEN: JCSAAZ; ISSN: 0590-9791
DT
     Journal ·
     Unavailable
LA
     cf. C.A. 50, 1617b. p-Cymene and CoF2 at 260-80° gives
AΒ
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perfluoro-(1-methyl-4-isopropyl cyclohexane) (I), b. 144-6°, nD16 1.3006. Similarly, p-ClC6H4Me or p-ClC6H4CF3 gives perfluoro (methylcyclohexane) (II), b. 75.5-5.8°, nD171.285, and 4-chlorotridecafluoro(methylcyclohexane) (III), h. 102-3°, nD17 1.314. II and PhMe passed through a tube at 540-55° give a compound b. below -10°, probably CF3CHFCF3, dibenzyl, and 4H-tridecafluoro-(methylcyclohexane) (IV), b. 85.1-6.3°. Similarly, II and Cl at 540-55° give CF3CFClCF3, b. 0°, and III. II and Br at 510° give CF3CFBrCF3, b. 15-16°, and 4-bromotridecafluoro(methylcyclohexane), b. 115.2-16.1°. Pyrolysis of II, perfluorodicyclohexyl, or perfluoro(isopropylcyclohexane) at 550-600° alone or mixed with one another gives only deep decomposition from which no pure compds. can be isolated. III and LiAlH4 give IV. IV refluxed 5 hrs. with aqueous KOH gives nonafluoro-4-trifluoromethyl-1cyclohexene (V), b. 75.4-5.9°, nD20 1.293 which adds Br to give the 1,2-dibromo compound, b. 156-9°, and Cl to give the 1,2-dichloro compound, b. 129-31°. V and KMnO4 give very hygroscopic perfluoro-(3-methyladipic) acid (VI), m. 60-1° [di-(Sbenzylthiuronium) salt, m. 208°; diethyl ester, b14 112°, nD12 1.361; diamide, m. 168-9°]. The di-K salt of VI decarboxylates when heated in ethylene glycol to give 1H,4H-heptafluoro-2trifluoromethylbutane, b. 63-5°. Heating this with KOH gives F ion.

ANSWER 9 OF 11 USPATFULL on STN L33 2003:192299 USPATFULL ANProcess for purifying octafluorocyclobutane, process for preparing the TIsame, and use thereof Horiba, Minako, Kanagawa, JAPAN INSuzuki, Yasuhiro, Kanagawa, JAPAN US 2003132099 A1 20030717 PIUS 2002-221443 A1 20020912 (10) ΑI WO 2002-JP148 20020111 20010115 PRAI JP 2001-6459 DTUtility FSAPPLICATION SUGHRUE MION, PLLC, 2100 PENNSYLVANIA AVENUE, N.W., WASHINGTON, DC, LREP 20037 Number of Claims: 32 CLMN

Exemplary Claim: 1 ECL

No Drawings DRWN

LN.CNT 1238

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A process for purifying octafluorocyclobutane according to the present ABinvention is characterized by contacting a crude octafluorocyclobutane containing impurities with an impurity decomposing agent under elevated temperature and then with an adsorbent to substantially remove the impurities from the crude octafluorocyclobutane.

According to the purification process or preparation process of octafluorocyclobutane of the present invention, the impurities such as fluorocarbon can be substantially removed and a high-purity octafluorocyclobutane can be easily obtained. The octafluorocyclobutane obtained by the purification process of the present invention is substantially free of impurities and therefore, can be used as an etching or cleaning gas for use in the production process of a semiconductor device or the like.

CAS INDEXING IS AVAILABLE FOR THIS PATENT

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ANSWER 10 OF 11 CAOLD COPYRIGHT 2004 ACS on STN
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CA63:16189b CAOLD AN

influence of substitution on CH.CF and CF.CF coupling consts. TI

Abraham, Raymond J.; Cavalli, L. AU

- TI tautomeric relations in S-containing heterocycles
- AU Wollner, Thomas E.
- L33 ANSWER 11 OF 11 CAOLD COPYRIGHT 2004 ACS on STN
- AN CA50:4048e CAOLD
- TI Reactions of highly fluorinated organic compds. (IX) 1H decafluoro 4 trifluoromethylcyclohexane, nonafluoro-4-trifluoromethyl-1-cyclohexene, and perfluoro-(3-methyladipic) acid
- AU Barlow, G. B.; Stacey, M.; Tatlow, J. C.

(FILE 'HOME' ENTERED AT 12:05:35 ON 03 AUG 2004)

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FILE 'REGISTRY' ENTERED AT 12:05:53 ON 03 AUG 2004
L1
              1 S 1,1,1,2,3,3,3-HEPTAFLUOROPROPANE/CN
L2
              1 S 2,2-DICHLORO-1,1,1,3,3,3-HEXAFLUOROPROPANE/CN
L3
              1 S 2-CHLORO-1,1,1,2,3,3,3-HEPTAFLUOROPROPANE/CN
L4
              1 S HYDROGEN FLUORIDE/CN
L5
             0 S 1,1,1,2,2,3,3,3-OCTAFLUOROPROPANE/CN
L6
             1 S OCTAFLUOROPROPANE/CN
L7
             1 S CHLORINE/CN
L8
              1 S HYDROGEN/CN
     FILE 'CAPLUS, USPATFULL, CA, CAOLD' ENTERED AT 12:13:49 ON 03 AUG 2004
L9
             22 S L1 AND L2
             20 S L9 AND L3
L10
            18 S L10 AND L4
L11
L12
             5 S L11 AND L7
             2 S L11 AND L8
L13
L14
             0 S L11 AND L6
             3 DUP REM L12 (2 DUPLICATES REMOVED)
L15
L16
             0 S L13 NOT L12
            17 S L9 NOT L12
L17
            12 DUP REM L17 (5 DUPLICATES REMOVED)
L18
L19
            58 S L1 AND L3
L20
            38 S L19 NOT L9
             9 S L20 AND L4
L21
             5 S L20 AND L8
L22
             4 DUP REM L22 (1 DUPLICATE REMOVED)
L23
L24
             9 S L21 NOT L22
L25
             5 DUP REM L21 (4 DUPLICATES REMOVED)
L26
             0 S L25 AND L8
L27
            29 S L20 NOT L21
L28
           24 S L27 NOT L22
L29
           13 DUP REM L28 (11 DUPLICATES REMOVED)
L30
             0 S L29 AND L8
L31
             0 S L29 AND L4
L32
             2 S L29 AND L7
         11 S L29 NOT L32
L33
L34
             0 S L33 AND L6
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